FORM I	ATTORNEY'S DOCKET NUMBER 10806-177 U.S. APPLIC, NO. (if known, see 37 CFR 1.5)							
	IATIONAL 00/02539	, APPLICATION NO.	INTERNATIONAL FILING DATE 16 March 2000	PRIORITY DATE CLAIMED 16 March 1999				
TITLE	F INVENT	TION MACRO	MOLECULAR COMPOUNDS					
APPLIC	ANT(S) FC	OR DO/EO/US HODD,	Kenneth A.; DILLINGHAM, Keith Alfred					
Applica	it herewith	submits to the United States Desig	nated/Elected Office (DO/EO/US) the following iter	ms and other information:				
1.	[X]	This is a FIRST submission of	items concerning a filing under 35 U.S.C. 371.					
2.		This is a SECOND or SUBSEC	QUENT submission of items concerning a filing und	ler 35 U.S.C. 371.				
3.	This express request to begin national examination procedures (35 U.S.C. 371(f)) at any time rather than delay examination until the expiration of the applicable time limit set in 35 U.S.C. 371(b) and PCT Articles 22 and 39(1).							
4.	[X]	A proper Demand for Internation	onal Preliminary Examination was made by the 19th	month from the earliest claimed priority date.				
5.	[X]	A copy of the International Application as filed (35 U.S.C. 371(c)(2)) a. [] is transmitted herewith (required only if not transmitted by the International Bureau). b. [X] has been transmitted by the International Bureau. c. [] is not required, as the application was filed in the United States Receiving Office (RO/US)						
6.	0	A translation of the Internationa	al Application into English (35 U.S.C. 371(c)(2)).					
7.	7.							
8.	D	A translation of the amendment	s to the claims under PCT Article 19 (35 U.S.C. 371	(c)(3)).				
9. 🕐	0	An oath or declaration of the in	ventor(s) (35 U.S.C. 371(c)(4)).					
10.	0	A translation of the annexes to	the International Preliminary Examination Report un	der PCT Article 36 (35 U.S.C. 371(c)(5)).				
Items 11	l. to 16. bel	ow concern other document(s) o	r information included:					
11.		An Information Disclosure Stat	ement under 37 CFR 1.97 and 1.98.					
12.		An assignment document for re	cording. A separate cover sheet in compliance with	37 CFR 3.28 and 3.31 is included.				
13.	[X] []	A FIRST preliminary amendments A SECOND or SUBSEQUENT						
14.	[]	A substitute specification.						
15.	1 3.	A change of power of attorney	and/or address letter.					
16.	[X] Other items or information: copy of published International Application No. WO 00/55212, including International Search Report							

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U.S. APPLIC. NO. (if known, see 37 CFR 1.50) INTERNATIONAL APPLICATION NO. PCT/EP00/02539				ATTORNEY'S DOCKET NUMBER 10806-177		
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17.	The following fees	are submitted:			CALCULATIONS	TTO OBE ONE
	Basic National Fee (37 CFR 1.492(a)(1)-(5)):					
[X]	Search Report has	been prepared by the EPO of	r JPO	\$860.00		
0	International prelin	ninary examination fee paid	to USPTO (37 CFR 1.482)	\$690.00		
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[] Neither international preliminary examination fee (37 CFR 1.482) nor international search fee (37 CFR 1.445(a)(2)) paid to USPTO \$1000.00						
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	endent Claims	1 -3 =	0	x \$80.00	\$	
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	<u>D. Kozlowski</u> D OR PRINTED N	AME		14 September 2001 DATE		

Docket No. 10806-177

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<u>PATENT</u>

IN THE UNITED STATES PATENT & TRADEMARK OFFICE

Applicant:

Kenneth A. Hodd et al

Paper No.:

Serial No.:

To be assigned

Group Art Unit:

Filing Date:

September 14, 2001

Examiner:

For: Macromolecular Compounds

PRELIMINARY AMENDMENT

Box PCT Assistant Commissioner for Patents Washington, DC 20231

Dear Sir:

Prior to calculation of the filing fee and first action by the Examiner, please amend the present application as follows:

In the Claims:

Please amend claims 10, 20, 21, 25, 27 and 28 to read as follows:

- 10. (Amended) Photocrosslinkers according to claim 1 provided with functional groups for crosslinking.
- 20. (Amended) A method of forming a macromolecular crosslinked network from a composition comprising a photocrosslinker according to claim 1 by irradiating said composition with light exceeding a wavelength of about 305 nm for a time sufficient to form a solid article.

- 21. (Amended) A method forming a macromolecular crosslinked network from a composition comprising a photocrosslinker according to claim 1 and at least one copolymerizable vinylic, acrylic or methacrylic monomer.
- 25. (Amended) A method according to claim 20, wherein an ophthalmic lens is produced.
- 27. (Amended) An ophthalmically acceptable composition comprising photocrosslinkers according to claim 1, having a refractive index greater than about 1.39 and a viscosity such that said composition can be injected through a standard cannula having a needle of 15 Gauge, or finer.
- 28. (Amended) A method for producing an intraocular lens, comprising injecting an ophthalmologically acceptable composition comprising photocrosslinker according to claim 1 into the capsular bag of the eye.

REMARKS

By the present Amendment, claims 10, 20, 21, 25, 27 and 28 are amended to omit their multiple dependency. Claim 28 is also amended to replace the recitation of a use with the recitation of a method for producing an intraocular lens, in accordance with the teachings of the specification. A Version With Markings Showing Changes Made is attached. It is believed that these changes do not involve any introduction of new matter, whereby entry is believed to be in order and is respectfully requested.

Respectfully submitted,

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VERSION WITH MARKINGS SHOWING CHANGES MADE

Claims 10, 20, 21, 25, 27 and 28 are amended as follows:

- 10. (Amended) Photocrosslinkers according to claim 1 [or 5] provided with functional groups for crosslinking.
- 20. (Amended) A method of forming a macromolecular crosslinked network from a composition comprising a photocrosslinker according to [any of claims 1 to 19] <u>claim 1</u> by irradiating said composition with light exceeding a wavelength of about 305 nm for a time sufficient to form a solid article.
- 21. (Amended) A method forming a macromolecular crosslinked network from a composition comprising a photocrosslinker according to [any of claims 1 to 11] <u>claim 1</u> and at least one copolymerizable vinylic, acrylic or methacrylic monomer.
- 25. (Amended) A method according to [any of claims 20 to 24] <u>claim 20</u>, wherein an ophthalmic lens is produced.
- 27. (Amended) An ophthalmically acceptable composition comprising photocrosslinkers according to [any of claims 1 to 19] <u>claim 1</u>, having a refractive index greater than about 1.39 and a viscosity such that said composition can be injected through <u>a</u> standard cannula having a needle of 15 Gauge, or finer.
- 28. (Amended) [The use of photocrosslinkers according to any of claims 1 to 19 in] A method for producing an intraocular lens, comprising injecting an ophthalmologically

acceptable composition <u>comprising photocrosslinker according to claim 1</u> [for injection] into the capsular bag of the eye.

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Macromolecular compounds

Field of invention

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The present invention relates to new photoinitiators capable of acting as photocrosslinkers providing a combination of photoinitiating and crosslinking processes.

Background of invention

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The UV curing of resin formulations is widely used in industry as the setting process for coatings, adhesives, and more recently paints. Such formulations may comprise a combination of vinyl, usually acrylate, monomers and crosslinkers, together with a photoinitiator. Other possible constituents of the formulations include crosslinkers and vehicles. In general an advantage of photocurable formulations is that the monomers act as their own vehicle, and the use of solvent is obviated, which has environmental advantages.

Advances in the technology of photocuring, improvements such as, those in UV lamps, cationic initiators for epoxide-based formulations, water borne coatings, and many novel monomers has enabled this production process to penetrate a number of important manufacturing sectors. Photopolymerization is now used in photoresists for printed circuits and microelectronics, for photolithography, magnetic recording media, glass-fiber laminates, and for medical devices, especially for dental and ophthalmic applications.

For the medical applications of photopolymerisation it is usual to employ visible light, rather than UV, to effect the cure of the resin formulation. The use of visible, usually blue, light avoids exposing patient and dentist or surgeon to harmful irradiation. Increasingly the merit of this approach is being recognized for industrial practice, where operatives also need protection from prolonged exposure to harmful UV.

European Patent 0800 657 describes a photoinitiator linked to a macromer structure which together with a copolymerizable monomer and a crosslinker is capable forming a polymerization product, such as an ophthalmic lens that retains photoinitiator radical in

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the resulting network. This is advantageous in medical applications wherein such potentially harmful radicals must be carefully controlled. However, this system would not be applicable for producing a polymerized product directly in the capsular bag in the eye since it is not directed to photoinitiators activated by light in the visible range. US Patent No. 4,536,265 discloses siloxane polyphotoinitiators to be used with a curable silicone resin. This system is UV curable and consequently it will not be applicable for photocuring in the living eye.

It is a characteristic of almost all, if not all, of the formulations used for aforementioned types of application that they are crosslinked. Crosslinking of the polymeric bases which constitute the coatings or artifacts of the aforementioned industrial products confers important advantages upon them. Crosslinked polymers have greater environmental (e.g. temperature and moisture) resistance, solvent resistance and dimensional and mechanical stability, than equivalent linear polymers. This is especially so for where the equivalent linear polymer are produced by photopolymerisation they have an atactic, non-crystalline, structure.

Crosslinking is introduced into photopolymerized products by including in the formulation for the resin, coating or gelling system, an acrylate, or similar, crosslinker, which is characterized by having two or more crosslinkable acrylate or vinyl functions. In some formulations this crosslinking species is a polymer of low molecular weight. The crosslinker copolymerizes with the monomers of the formulation to produce a network structure.

It is the object of the present invention provide compounds which act as photocrosslinkers for vinyl, acrylate and methacrylate monomers and acrylated silicone compositions, especially in solution.

It is also an important object of the present invention to provide photocrosslinkers with capability to act in aqueous solutions, especially on water soluble macromolecular particles having functional groups for crosslinking.

It is another object of the present invention to provide photocrosslinkers with enhanced photoactivity (100 % conversion of monomer to polymer in aqueous solution) which reduces photoinitiztor residues to a minimum, especially, vinyl modification of

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photoinitiator component and thereby reducing compositional drift, Draize and other environmental hazards.

The invention as presented below will explain how the mentioned objects are met while discussing further obvious advantages.

Description of the invention

The present invention pertains to macromolecular hydrophilic photocrosslinkers having a general formula $(A)_n(B)_m(C)_p$, wherein

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- (i) A, B and C are units of substituted ethylene or siloxane groups in the macromolecular structure;
- (ii) A, B and C are randomly distributed and the unit C carries a photoactive group;
- (iii) n = 0.98 mole %, m = 0.98 mole %, n+m = 50.98 mole % and p = 0.5.50 mole %.

When the photoactive groups of units C are exposed to light of determined wavelengths above 305 nm, radicals are generated which are retained on the macromolecular photocrosslinkers and will react to form a crosslinked network structure. Preferably the final structure is solid article.

The photocrosslinker further preferably further comprises functional groups for crosslinking. Such groups are conventionally vinylic, acrylic or methacrylic groups and their nature and introduction on polymeric backbone are well known to persons skilled in the art and will be referred to as "functional groups for crosslinking".

According to one aspect of the invention a fluid composition of the photocrosslinker in a suitable amount can be directly crosslinked into the final solid product upon sufficient irradiation. In another aspect the composition for crosslinking into a solid article comprises suitable amounts of the photocrosslinker and a polymer carrying functional groups for crosslinking. The photocrosslinker in such a system will thereby replace the conventional combination of crosslinker and photoinitiator.

Applicable polymers with suitable functional can readily be provided with the skilled person for the purpose of crosslinking desired articles. For example it would be

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conceivable to employ polymers having a sufficiently high refractive index to be acceptable as intraocular lenses. Suitable polymers can be, for example, be found in International Patent Application PCT/EP99/07718. In a still another aspect of the present invention, the photocrosslinkers can be employed in a composition, preferably an aqueous composition further comprising at least one copolymerizable vinylic, acrylic or methacrylic monomer. Such monomers and combinations thereof are well known in the art and will not be described herein in further detail. It is, however, to be understood that the photocrosslinker will replace conventional crosslinking agents and their combination with photoinitiators in such systems.

It is highly preferred that the photoactive groups of the photocrosslinkers comprise a phosphine oxide, in order to generate the necessary radicals for crosslinking from the exposure of visible light. More preferably, the photoactive group is an acyl- or aroyl phosphine oxide.

According to a preferred aspect, the photoactive group is linked to the ethylene groups of units C of the photocrosslinkers by a linking group comprising a phenylene group. Optionally, such a phenylene group is substituted in order to obtain more stability.

According to one embodiment of the invention, the photocrosslinkers comprises substituted ethylene units A, B, C of a macromolecular photocrosslinker in according to:

20 A=-CH₂-C($\mathbb{R}^1\mathbb{R}^2$)-, B=-CH₂-C($\mathbb{R}^1\mathbb{R}^3$)-, C=-CH₂-C($\mathbb{R}^1\mathbb{R}^4$)-, wherein

R¹ is hydrogen or methyl;

R² is -CON(Me)₂, -CO₂CH₂CH₂OH, -OCOCH₃, -OCOCH₂CH₂Ph, -OH or a lactam group;

 R^3 is $-CON(Me)_2$, $-CO_2CH_2CH_2OH$, $-OCOCH_3$, $-OCOCH_2CH_2Ph$, -OH or a lactam group when B is $-CH_2-C(R^1R^3)$ - with the proviso that R^2 and R^3 are not the same; and R^4 is $-R^5C(O)P(O) R^6R^7$ or $-R^5P(O)R^6OC(O)R^7$, wherein R^5 , R^6 and R^7 are selected among same or different aryl groups comprising phenyl, methylphenyl, dimethylphenyl,

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trimethylphenyl, methoxyphenyl, dimethoxyphenyl, trimethoxyphenyl, methylolphenyl, dimethylolphenyl, trimethylolphenyl or styryl radicals, or

In the general formula above, -OH denotes a hydroxyl group, Me a methyl group and Ph is a phenyl group. The lactam group typically is a heterocyclic ring structure of 4 to 7 atoms of which at least one is nitrogen. A suitable such lactam group provides a N-vinyl-pyrrolidone structure as one of units A or B on said ethylenic backbone. It is also to be understood that besides the mentioned substituents functional groups for crosslinking can be added to the macromolecule in accordance with conventional methods.

In one advantageous aspect of this embodiment, the photocrosslinkers, R² and R³ according to above are selected so as to form a water-soluble molecule.

Suitable units A and B in the general formula (A)_n(B)_m(C)_p are selected among, but not limited to, N-vinylpyrrolidone (NVP), 2-hydroxyethylmethacrylate, N-N-dimethylacrylamide and vinyl acetate. The vinyl acetate referred to preferably will be hydrolyzed conventionally to vinyl alcohol. It is also referred to Table 1 below in the exemplifying part of the description for a number of specific photocrosslinkers based on such units (or co-monomers) and 4-vinylbenzoyl-diphenylphosphine oxide (VBPO) as a photoinitiating group. Accordingly, VBPO units constitute units C in said general formula above. Some especially suitable water soluble, blue light activated photocrosslinkers according to the present invention comprise NVP together with vinyl acetate units, N,N-dimethylacrylamide units alone or together with 2-hydroxyethylethacrylate units, all combined with VBPO units. These photocrosslinkers demonstrate high conversion rate (monomer to polymer) and suitably high stability in aqueous solution. This type of photocrosslinkers can be prepared by conventional radical polymerization.

According to another embodiment, the photocrosslinkers described above with general formula can comprise units A, B and C which are siloxane monomer units having a formula

 $-R_aR_bSiO$ -, wherein R_a and R_b in units A and B are selected among lower substituted or unsubstituted alkyl groups, aryl groups and arylalkyl groups. Preferably, at least on of R_a and R_b is an aryl or arylalkylgroup. More preferably R_a and R_b is substituted with one or more fluorine atoms. Alkyl groups in this context means a C_1 to C_{10} alkyl group which is

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straight or branched.

According to a preferred aspect of this embodiment the siloxane units comprising substituents in accordance with:

A is $-Si(R^1R^2)$ -O-, B is $-Si(R^1R^3)$ -O- and C is $-Si(R^1R^4)$ -O-, wherein

R¹ is C1 to C6 alkyl; R² is C1 to C6 alkyl or phenyl; R³ is R¹, R² or C1 to C6 fluroalkyl;

R⁴ is -R⁵R⁶C(O)P(O) R⁷R⁸ or -R⁵R⁶P(O)R⁷OC(O)R⁸, wherein R⁵ is a spacing group; R⁶, R⁷ and R⁸ are selected among same or different aryl groups comprising phenyl, methylphenyl, dimethylphenyl, trimethylphenyl, methoxyphenyl, dimethoxyphenyl, trimethylolphenyl, trimethylolphenyl or styryl radicals.

The aliphatic spacing group R⁵ is preferably comprises between one and ten atoms and suitably

The spacing group is $(-CH_2)_n$, wherein n is between 1 and 10.

According to an aspect of the invnetion particularly suitable for the production of ophthalmic lenses, the photocrosslinkers has radicals connected to the polysiloxane backbone such that R¹ is methyl; R² is methyl or phenyl; and R³ is R¹, R² or -CH₂CH₂CF₃. Such polysiloxane photocrosslinkers may have functional acrylic groups in its terminal ends. Polysiloxanes of this type and their applicability and advantages, especially for injectable intraocular lenses, are disclosed in the International Patent Application PCT/EP99/07781 which document herewith is incorporated as a reference.

The present invention further involves a method of forming a macromolecular crosslinked network from a fluid composition comprising photocrosslinkers according to any of the mentioned embodiments by irradiating said composition with light exceeding a wavelength of about 305 nm for a time sufficient to form a solid article. The composition can comprise said

50 photocrosslinkers at least one copolymerizable vinylic, acrylic or methacrylic monomer.

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or the composition can comprise a polymer provided with functional vinylic, acrylic or methacrylic groups. It would be obvious to the skilled person to combine any such monomers and polymers together with the inventive photocrosslinkers and also, if found advantageous, combining the composition with a conventional crosslinker suitable for the specifically selected composition. It is further to be understood that the constituents of such a composition shall be selected so as be sufficiently compatible to each other and the selected fluid environment, for example depending on if photocrosslinkers having ethylene or polysiloxane backbone are selected.

In an especially advantageous application of the method, a medical device or medical implant, such an ophthalmic lens is produced by means of a conventional molding method wherein the photocrosslinking into a network is a conventional curing process. The inventive method is particularly suitable for producing an intraocular lens by means of injection and subsequent photocrosslinking direct in the capsular bag of eye, from which the natural lens has been surgically removed.

It is also a part of the present invnetion provide an ophthalmically acceptable composition comprising the new photocrosslinkers. Such a composition will typically have a refractive index greater than about 1.39 and a viscosity such that said composition can be injected through standard cannula having a needle of 15 Gauge, or finer. Such a composition can further comprise of any suitable constituents as outlined above that can be a part of the network provided by the subsequent photocrosslinking.

The photocrosslinkers according to the present invention provide for a combination of photoinitiating and crosslinking processes. It is an important feature of the present invention to effect this combination of function by attaching photoactive groups to a polymeric or macromolecular structure. The photoactive groups, when exposed to light of the appropriate wavelength, will undergo photoinduced scission and generating radicals, which are retained on the polymeric or macromolecular structure. These retained radicals then initiate, terminate, or, in some other way participate in the gel forming process that is the objective of the radiation cure of the photomaterial. The use of the inventive photocrosslinkers confers distinct advantages, both chemical and environmental, as compared with the combination of a separate photoinitiator and crosslinker. In a chemical

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context the use of a photocrosslinker gives opportunities to produce networks that are more homogeneous than those produced by photocuring conventional photocurable systems. The latter systems, involving as they do, combinations of monomers, have structures dependent on the reactivity ratios of the monomers and crosslinkers. Often, for example, in a coating being manufactured at high rates of production, a crosslinker is selected because of its high reactivity. Disparities in the reactivates of the components of a formulation gives rise to compositional drift, the change of the average unit composition during the course of a polymerization, and this in relation to a reactive crosslinker implies that sections of a network forming later, in the curing process, have a lower crosslink density than sections formed earlier. Improving the homogeneity of crosslinked networks is a subject receiving greater attention as the technical demands imposed on industrial products increases. Homogenous networks have, for example, higher fracture toughness and better optical properties heterogeneous networks. The shrinkage occurring during their formation is more uniform allowing for more precision in castings. The benefits of using a photocrosslinker as a network former, as compared with a combination of photoinitiator and crosslinker, arise because the radical species they produce act as crosslinkers via the polymer chain to which they are attached. Further such radicals are generated throughout the setting phase, their concentration being controlled by the photoinitiating species' quantum efficiency and the intensity of the light, which may be modulated during the setting, in addition to its concentration. This distinction results in the formation of networks having a more controlled and homogeneous structure.

Retaining photoinitiator residues in the network of a medical product, such as a contact lens or a dental filling has desirable physiological implications. Further photocrosslinkers because of their polymeric, or macromolecular, nature are more acceptable, environmentally, than many conventional crosslinkers which are known to cause skin and lung irritation.

Within the context of the present invention, it is possible to substitute a photocrosslinker, either completely, or partially, for a combination of a conventional photoinitiator and a conventional crosslinker. Alternatively, the inventive

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photocrosslinkers can be used in combination with a conventional photoinitiator or a conventional crosslinker, as will be understood by practitioners skilled in formulating systems for crosslinking.

Persons skilled in this art will also appreciate that the inventive photocrosslinkers as described herein for photoactive systems responsive to visible light may be applied equally to systems responsive to UV light, so the present invention is of very general applicability.

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Detailed and exemplifying description of the invention

Example 1

5 PHOTOCROSSLINKER POLYMER PREPARATIONS

Table 1

photocrosslink	VBPO	Comonomer	Comonomer
ers	(mole%)	1 (mole%)	2 (mole%)
	1		_
P31-1	3.5	HEMA(5)	NVP(91.5)
P32-1	3.5	VAc(10)	NVP(86.5)
P40-3	4	DMA(96)	none
P40-4	4	PEMA(96)	none
P41-1	6	DMA(94)	none

The following Examples describe the preparation of P32-1(3). P40-3 & P41-1 (comparison), and P40-4 respectively. In addition examples demonstrating photocrosslinkers of DMA and 4-vinyl-2,6-dimethylcbenzoylphosphine oxide are added (Examples 1E and 1F).

15 Example 1A

Photocrosslinker Copolymer employing N-Vinylpyrrolidone and Vinyl acetate

This preparation, on an 8g monomer scale, used monomers in the molar ratio:
86.5 parts N-vinylpyrrolidone (VP): 10 parts vinyl acetate (Vac): 3.5 parts
vinylbenzoyldiphenylphosphine oxide (VBPO).

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Methoxydiphenylphosphine, 0.520g, was weighed to a dried 100ml twin-neck flask, with one neck septum sealed, and coated in aluminium foil to exclude light. Toluene, 3 ml, and a magnetic stir bar were added and the flask flushed with dry nitrogen. The stopcock was briefly removed and 4-vinylbenzoyl chloride, 0.409g, added, the flask being again flushed with dry nitrogen, then placed in a bath at 65°C, with magnetic stirring.

After 15 minutes, the other monomers: VP, 6.620g, and Vac, 0.595g, were diluted with a previously prepared solution of azobisisobutyronitrile (AIBN), 0.080g in 8ml toluene, and the mixture injected to the flask and rinsed in with a further 4ml toluene. The polymerization mixture was heated at 65°C with magnetic stirring for 8 hours, yielding a clear pale yellow solution, which was precipitated, in subdued light, to diethyl ether. The supernatant was discarded and the pale sludge-like precipitate taken up in 30ml methanol and reprecipitated to ether as a curdy precipitate. The supernatant was decanted, and the polymer product dried to constant weight under vacuum at 35°C. Yield was 5.751g (72%) of friable pale yellow polymer. Elemental analysis gave 0.65% P, corresponding to 6.9%ww VBPO units (0.209mmol/g), and 10.70% N, corresponding to 84.5%ww VP units, and thus a mean unit mass of 115 Daltons. SEC gave Mn 32,000, Mw 103,000. This implies a number average chain length of ca.280 units, with ca.7 photoactive units per chain.

Example 1B

Photocrosslinker Copolymer employing N-Dirnethylacrylamide (I)

25 In this example, 4-vinylbenzoyldiphenylphosphine oxide (VBPO), 4 mol%, was copolymerized with N,N-dimethylacrylamide (DMA), 96 mol%, on a 6g scale.

Methoxydiphenylphosphine, 0.481g, was weighed to a dried 24x150mm Quickfit tube, and 2.5ml dry toluene added. The tube was then wrapped in aluminium foil to exclude light. 4-Vinylbenzoyl chloride, 0.368g, and a magnetic stir bar were added, and the tube septum sealed, N₂ flushed, and placed in a bath at 65°C with stirring. After 15 minutes a solution of DMA, 5.26g, and AIBN, 0.060g, in toluene, 5ml, was injected by

#II

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syringe and rinsed in with a further 3.6ml toluene. The mixture was stirred 6h at 65°C, yielding a viscous orange-yellow solution, which was diluted with methanol and precipitated in diethyl ether. The product was reprecipitated from methanol to ether, and vacuum dessicated at room temperature. Yield, 2.56g (43%) of friable pale yellow polymer, analysis 0.82% P corresponding to 8.8%ww VBPO units (0.265mmol/g). GPC using poly(ethylene glycol) standards gave Mn 25,000; Mw 100,000.

Example 1C

10 Photocrosslinker Copolymer employing N-Dimethylacrylamide (II)

Example 2B was repeated on a 12 g scale, but with monomer ratio 6 mol% VBPO (2.12 g), 94 mol% DMA (9.89 g), with 0.120 g AIBN, 22.3 ml toluene, and polymerization time extended to 8h at 65°C. The yield was 7.17g (60%) of yellow polymer, analysis 1.49% P corresponding to 16.0 %ww VBPO (0.481mmol/g).GPC gave Mn 12 000 and Mw 88000.

Example 1D

20 Photocrosslinker Copolymer employing 2-Phenylethyl methacrylate

In this example, 4-vinylbenzoyldiphenylphosphine oxide (VBPO), 4 mol%, was copolymerized with 2-phenylethyl methacrylate (PEMA), 96 mol%, on a 6g scale.

Methoxydiphenylphosphine, 0.271g, was weighed to a dried 24x150mm Quickfit tube, and 2.5ml dry toluene added. The tube was then wrapped in aluminium foil to exclude light. 4-Vinylbenzoyl chloride, 0.204g, and a magnetic stir bar were added, and the tube septum sealed, N₂ flushed, and placed in a bath at 65°C with stirring. After 15 minutes a solution of PEMA, 5.60g, and AIBN, 0.060g, in toluene, 5ml, was injected by syringe and rinsed in with a further 3.6ml toluene. The mixture was stirred 6h at 65°C, yielding a fairly viscous pale yellow solution, which was diluted with chloroform and precipitated to methanol. The product was reprecipitated from chloroform (with THF

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added to clarify the solution), and vacuum dessicated at room temperature. Yield, 4.67g (78%) of friable pale yellow polymer, analysis 0.48% P corresponding to 5.2%ww VBPO units (0.155mmol/g). GPC in THF using polystyrene standards gave Mn 49,300; Mw 108,500.

Example 1E

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In this example, 4-vinyl-2,6-dimethylbenzoyldiphenylphosphine oxide (VDMBPO), 4 mol%, was copolymerized with N,N-dimethylacrylamide (DMA), 96 mol%, on a 12g scale.

Methoxydiphonylphosphine, 0.979g, was weighed to a dried flask and 5ml dry toluene added. The flask was wrapped in aluminium foil to exclude light. 4-Viny-2,6-dimethyllbenzoyl chloride, 0.885g, and a magnetic stir bar were added, and the flask septum sealed, N₂ flushed, and placed in a bath at 65°C with stirring. After 15 minutes a solution of DMA, 10.426g, and AIBN, 0.121g, in toluene, 9.3ml, was injected by syringe and rinsed in with a further 8ml toluene. The mixture was stirred 8h at 65°C, yielding a viscous pale yellow solution, which was diluted with 20ml ethanol and precipitated in diethyl ether. The product was reprecipitated from ethanol to hexane, and vacuum dessicated at room temperature. Yield, 8.53g (71%) of friable pale yellow polymer, analysis 0.58% P corresponding to 6.75%ww (1.95mol%) VDMBPO units (0.187meq/g).

The polymer was water soluble and showed excellent hydrolytic stability; tested over the course of a year the product showed no measurable decrease in photoactivity.

GPC gave Mn 6,000; Mw 26,000.

25 Example 1F

Example 1E was repeated employing VDMBPO 5 mol and DMA 95 mol%. Yield was 43% of pale yellow polymer, analysis 0.86% P corresponding to 10.0%ww (2.97mol%) VDMBPO units (0.278meq/g). GPC gave Mn 7,000; Mw 32,500.

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Example 1G

Example 1F was repeated employing VDMBPO 5 mol% and DMA 95 mol%. Yield was 55% of pale yellow polymer, analysis 0.73% P corresponding to 8.5%ww (2.49mol%) VDMBPO units (0.236meq/g). GPC gave Mn 5,600; Mw 24,000.

EXAMPLE 1H

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1,3,5-trimethylbenzoyl-styrylphenylphosphine oxide (TMBSPO), 4 mol%, was copolymerized with N,N-dimethylacrylamide (DMA), 96 mol%, on a 12g scale.

First methoxystyrylphenylphosphine, 0.800g, was weighed to a dried flask and 5ml dry toluene added. The flask was wrapped in aluminium foil to exclude light. 1,3,5-trimethyllbenzoyl chloride, 1.061g, and a magnetic stir bar were added, and the flask septum sealed, N₂ flushed, and placed in a bath at 65°C with stirring. After 15 minutes subsequently a solution of DMA, 10.241g in 15 mL of toluene, and AIBN, 0.120g in 5.0 mL of toluene, were injected by syringe. The mixture was stirred 8h at 65°C, yielding a viscous pale yellow solution, which was diluted with 20ml ethanol and precipitated in diethyl ether. The product was reprecipitated from ethanol to diethylether, and vacuum dessicated at room temperature. Yield was 55% of pale yellow polymer, analysis 0.87% P corresponding to 10.4%ww (2.40mol%) TMBSPO units (0.227meq/g). GPC gave Mn 9,000; Mw 35,000.

The experiment was repeated employing TMBSPO 2.5 mol% and DMA 97.5 mol%. Yield was 79% of pale yellow polymer, analysis 0.43% P corresponding to 5.1%ww (1.19mol%) TMBSPO units (0.112meq/g). GPC gave Mn 15,000; Mw 94,000.

Finally, the experiment was repeated employing TMBSPO 4 mol% and PEMA 96 mol%. Yield was 68% of friable pale yellow polymer, analysis 0.56% P corresponding to 5.4%ww TMBSPO units (0.149mmol/g). GPC gave Mn 19,000 and Mw 165,000.

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Example 2

The following examples refer to photopolymerization including the inventive photocrosslinkers compared with photopolymerization with commercially available photoinitiators.

Example 2A

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The state of the art photoinitiator Irgacure 1800 (ex Ciba-Geigy, 10.0 mg) was dissolved, in subdued light, in 2-hydroxyethylmethacrylate (HEMA, ophthalmic grade ex. Polysciences, 970 mg) and 1,6-dihydroxyhexane diacrylate (HDDA, ex, 20.0 mg), and a 10.0 mg sample pipetted into an open DSC aluminum sample pan. The sample pan, covered with a cover-slip of thin glass, was placed in the sample position of the head of a TA Instruments Differential Photocalorimeter (DPC). The temperature of the head was allowed to stabilize under N₂ at 37°C (or in some cases 23°, and the sample irradiated with blue light at an intensity of 8-9 mWcm⁻².

The area of the polymerization exotherm was determined by conventional computation and the Jg-1 of monomer calculated. From the Jg-1 the percentage conversion of monomer to polymer was calculated using a literature value for the latent heat of polymerization of the monomer, ΔH_p . The findings are collected in Table 2.

Using the same composition as was used for the DPC tests discs (2mm thick x 16mm diameter) of polyHEMA were cast in PTFE casting cells. About 500mg of the mixture of monomers and photoinitiator were introduced into the cell which was closed with a glass slide and irradiated with blue light, either from a blue light dental gun, or from a proprietary light generator (Efos Novacure), for 3min.

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Example 2B

The method described in Example 2A was repeated using with the state of the art photoinitiator Lucirin TPO (ex BASF, 10.0mg) instead of Irgacure 1800.

Example 2C

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The method described in Example 2A was repeated using HEMA (900.0mg), no HDDA, and, instead of Irgacure 1800, a photocrosslinker according to the present invention (P31-1, see Table 1. for composition, 100.0mg)

Example 2D

The method described in Example 2C was repeated using a photocrosslinker according to the present invention (P32-1, see Table 1. for composition, 100.0mg).

Example 2E

The method described in Example 2C was repeated using a photocrosslinker according to the present invention (P40-3, see Table 1. for composition, 100.0mg).

Example 2F

The method described in Example 2C was repeated using a photocrosslinker according to the present invention (P41-1, see Table 1. for composition, 100.0mg).

Example 2G

The method described in Example 2A was repeated using a photocrosslinker according to the present invention (P32-1, 100.0mg) instead of Irgacure 1800, HEMA (600.0mg),

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water(300mg) and no HDDA.

Example 2H

As Example 2G, using P40-3 (50.0mg) to replace P32-1, and HEMA (500.0mg), water (450.0mg).

Example 2I

10 As Example 2H using P41-1(50.0mg) to replace P40-3.

Example 2.J

The method described in Example 2A was repeated using 2-phenylethylacrylate (PEA, 990.0mg, ex Polymer & Dajac Laboratories) instead of HEMA and no HDDA.

Example 2K

The method described in Example 2A was repeated using instead of Irgacure 1800 a photocrosslinker (P40-4, see Table 1. for composition, 100.0mg) and PEA (900mg) but no HDDA or HEMA.

The % conversions of monomer to polymer in Table 2., Examples 2A and 2B, the commercial photoinitiators, and the photocrosslinkers, Examples 2C to 2E, are comparable showing that the photocrosslinkers behave as efficient photoinitiators, especially giving regard to the concentrations of photoactive species, the acylphosphine oxide (shown in Table 1) Further when these findings are compared with Examples 2G to 2I the comparison reveals that correctly designed photocrosslinkers (Examples 2H and 2I) exhibit 100% conversions in solution in water.

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For the 2-phenylethylacrylate monomer the photocrosslinker P40-4, based on 2phenylethylmethacrylate, is also very efficient as a photoinitiator (comparing Examples 2J and 2K) giving 100% conversion of monomer to polymer gel, as judged from the heat of polymerization (based on experimentally determined ΔH_p).

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Table 2. A comparison of the completeness of blue light photopolymerisation of HEMA, HEMA in water, & PEA using low molecular weight photoinitiators and photocrosslinkers

Ex.	Formulation ²	Heat of Polym.	Polym.	Conversion
No.	(wt%)[m.eq photoactive	(Jg ⁻¹)	time (min)	%
	ingredient ^b /100g]			
2A	HEMA(97)HDDA(2)I1800(1)[0.51	351	3.5	80
]			
2B	HEMA(97)HDDA(2)TPO(1)[2.9]	357	1.5	82
2C	HEMA(90)P31-1(10)[2.0]	308	6	70
2D	HEMA(90)P32-1(10)[2.3]	309	3	71
2E	HEMA(90)P40-3(10)[2.7]	307	2	70
2F	HEMA(90)P41-1(10)[4.8]	361	1.5	8 2
2G	HEMA(60)H ₂ O(30)P32-1(10)[2.3]	>275	>7	>63
2H	HEMA(50)H ₂ O(45)P40-3(5)[1.4]	452	7	100(approx.)
21	HEMA(50)H ₂ O(45)P41-1(5)[2.4]	454	6	100(approx.)
2Ј	PEA(99)I1800(1)[0.51]	455	2.5	100(арргох.)
2K	PEA(90)P40-4(10)[1.6]	456	3.5	100(approx.)

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^aPhotocrosslinkers, and monomer HEMA, as Table 1: commercial photoinitiators

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I1800, bis(2,6-dimethoxybenzoyl)-trimethylpentylphosphine oxide (25%) + 1-hydroxy-cyclohexylphenylketone (75%)(Irgacure 1800 ex Ciba-Geigy)
TPO, 1,3,5-trimethylbenzoyldiphenylphosphine oxide (Lucirin TPO ex BASF):
monomer PEA, 2-phenylethylacrylate: crosslinker HDDA, hexan-1,6-diol diacrylate
bm.eq. of acylphosphine oxide/100g of formulation.

Example 3

Examples for Gelation Tests:

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Examples 3A and 3B

Using the formulations described above in Examples 2J and 2K and the casting method described in Example 2A discs were prepared.

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Example 3C

Irgacure 2959 (ex Ciba-Geigy, 10.0 mg) was dissolved, in subdued lighting, in 2-hydroxyethylmethacrylate (HEMA, ophthalmic grade ex Polysciences, 550.0 mg) and water (440.0 mg). Test discs (2mm thick x 16mm diameter) of polymer were cast in PTFE casting cells. About 800mg of the mixture of monomers and photoinitiator were introduced into the cell which was closed with a glass slide and irradiated with light from a proprietary light generator (Efos Novacure), for 3 min.

25 Example 3D

As Example 3C with Irgacure 2959 (30.0 mg), HEMA (540.0 mg) and water (430.0 mg).

Example 3E

30 As Example 3C with P40-3 (100.0 mg) replacing Irgacure 2959, HEMA (500.0 mg), and

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water (400.0mg).

Example 3F

As Example 3C with P41-1 (70.0mg) replacing Irgacure 2959, HEMA (510.0mg), and water (420.0mg).

Example 3G

As Example 4C with P40-4 (50.0mg) replacing Irgacure 1800, PEA (900.0mg), and additional crosslinker, CE7-2 (2-phenylethylmethacrylate/2-hydroxy-3-acryloxypropylmethacrylate copolymer [0.9:0.1 mole ratio], 50.0mg).

Example 3H

As Example 3G with Irgacure 1800 (21.0mg) replacing P40-4, PEA (940.0mg), and crosslinker, CE7-2 (2-phenylethylmethacrylate/2-hydroxy-3-acryloxypropylmethacrylate copolymer [0.9:0.1 mole ratio], 60.0mg).

20 Example 3I

As Example 3B with PEA (750.0mg), and photocrosslinker, P40-4 (250.0mg).

In Table 3, are collected the tests made to check the gelation of the different formulations.

Where a composition is gelled it does not dissolve in solvent, but swells to an extent related to its crosslink density. Uncrosslinked (sol) polymers dissolve.

Examples of monomers photopolymerized with conventional photoinitiators of low molecular weight, nos. 4A, 4C and 4D dissolved readily in the appropriate solvent, water for polyHEMA, and acetone for polyPEA. Example no. 4B showed an intermediate

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behavior and dissolved partially in acetone leaving some residual gel. Increasing the proportion of photocrosslinker to 25% (3.9m.eq. of acylphosphine oxide, Example 4I or, adding separate crosslinker, CE7-2 (Example 4G, see below) produced acetone insoluble gel.

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CE7-2, a polyPEMA which is unsaturated and PEA miscible, being a copolymer of 2phenylethylmethacrylate/2-hydroxy-3-acryloxypropylmethacrylate [0.9:0.1 mole ratio], was employed as a supplementary crosslinker to the photocrosslinker P40-4, in Examples 4G and 4H. That CE7-2 is an effective cross-linker for photopolymerized PEA is demonstrated in example no. 4H, where in combination with Irgacure 1800 it also yields a gelled product upon irradiation. The products upon irradiation are transparent gelled elastomers of high refractive index (RI>1.54), similar in properties to PEA/PEMA copolymers.

Examples 3E and 3F which used photocrosslinkers to replace conventional 15 photoinitiators for HEMA/water compositions were gelled and did not dissolve in water, unlike examples 4D and 4E.

Table 3. Gelation tests on photopolymerized materials, shewing effect of photocrosslinkers

Ex. No.	Formulation (wt%) ¹	Effect of Solvent	Comments
3A	PEA(99)I1800(1)	Dissolves in Acetone	Not Crosslinked
3B	PEA(90)P40-4(10)	Dissolves & Swells in Acetone	Lightly Crosslinked
3C	HEMA(55)H ₂ O(44)I2959 ² (1)	Dissolves in Water	Not Crosslinked
3D	HEMA(54)H ₂ O(43)I2959(3)	Dissolves in Water	Not Crosslinked
3E	HEMA(50)H ₂ O(40)P40-3(10)	Swells in Water	Crosslinked Gel

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3F	HEMA(51)H ₂ O(42)P41-1(7)	Swells in Water	Crosslinked Gel
3G	PEA(90)CE7-2(5)P40-4(5)	Swells in Acetone	Crosslinked Gel
3H	PEA(94)CE7-2(6)I1800(2.1)	Swells in Acetone	Crosslinked Gel
31	PEA(75)P40-4(25)	Swells in Acetone	Crosslinked Gel

¹ See Tables 1. & 2., and text for an explanation of materials codes ²I2959, 2-hydroxy-4'-hydroxyethoxy-2-propiophenone (UV curing)

5 The crosslinked structure of the water swollen hydrogels (4E and 4F) was confirmed by stress relaxation tests.

Example 4

10 The method described in Example 2A was repeated using the following formulations:

Formulations (wt %)

- 4A. water (80)/photocrosslinker according Example 1F(20)
- 4B. water (80)/photocrosslinker according Example 1H(20)
- 15 4C. HEMA(45)/water(35)/photocrosslinker according to Example 1C(20)
 - 4D. HEMA(45)/water(35)/photocrosslinker according to Example 1F(20)
 - 4E. HEMA(45)/water(35)/photocrosslinker according Example 1H(20)
 - 4F. HEMA(45)/H20(35)
- The coherent and clear gels resulted from the irradiation of the formulations with blue light, and their relative crosslinked nature was characterized in two ways. The first method was to measure the stress relaxation of the networks, using a Rheometrics RDA-11, and the second method used was to measure the smiling of the gels in water.
- 25 Stress Relaxation Tests-Method

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The RDA-11 was set up with the 16mm gelled sample damped between parallel plates of 25 mm, heated to 35°C, and a strain of 30% applied. During the test the instrument measures the instantaneous stress necessary to maintain 35%, and plots the instantaneous shear modulus (Gi) against log t In Table 3, the percentage reductions in the modulus Gi for the formulations 4A to 4F between i = 10 and 100 s are compared as (G(10) -G(100)/G(10))x100 both before and after swelling in water. The results confirm that the photocrosslinked gel possess coherent network structures.

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Table 3. Average Stress Relaxations of Photocrosslinked Formulations 4A to 4F, measured at 35°C

Average stress relaxation	4A	4B	4C	4D	4E	4F
Before swelling- disc1	No result		4.9	3.4	No result	Not measurabl e
Before swelling- disc2	9.3		15.5	10.9	29.4	
After swelling- disc1			21.4	17.4		
After swelling- disc2	19.1		19.0	13.0	12.4	Not measurabl e

Swelling Test Method 15

Samples discs from formulations 4A through 4F were weighed, immersed in water for 24 hours at 20°C, dried, and reweighed. Table 4 compares the water absorbed by each formulation on a percentage basis.

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Table 4.

Percentage water absorbed at 25°C by photocrosslinked gels

Water absorbed (wt %)	4A	4B	4C	4D	4E	4F
Discl			222	191		Not measurabl e
Disc2	130	219	230	172	255	Not measurabl e

It was observed that upon irradiation with blue light, the formulation prepared without a crosslinker (4F) did not gel and that no discs suitable for any measurements were formed. Satisfactory discs were prepared from other formulations and stress relaxation results and the water absorption results were in agreement with the sequence: most highly crosslinked 4A>4C<4E least highly crosslinked.

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Claims

- 1. Macromolecular photocrosslinkers having a general formula
- 5 $(A)_n(B)_m(C)_p$, wherein
 - (i) A, B and C are units of substituted ethylene or siloxane groups in the macromolecular structure;
 - (ii) C carries a photoactive groups;
- 10 (iii) n = 0.98 mole %, m = 0.98 mole %, n+m = 50.98 mole % and p = 0.5.50 mole %;

and when said photoactive groups are exposed to light of determined wavelengths above 305 nm, radicals are generated and retained on the macromolecular photocrosslinkers and reacting so as to accomplish a crosslinked network structure.

- 2. Photocrosslinkers according to claim 1 characterized in that said photoactive group comprises a phosphine oxide.
- 3. Photocrosslinkers according to claim 2 characterized in that the photoactive group is an acyl- or aroyl phosphine oxide.
 - 4. Photocrosslinkers according to claim 3 characterized in that the photoactive group is linked to the ethylene groups of units C by a linking group comprising a phenylene group, said phenylene group being optionally substituted.
 - 5. Photocrosslinkers according to claim 1, wherein the ethylene units A. B, C of the macromolecular structure comprises substituents in accordance with:

$$A = -CH_2 - C(R^1R^2) - , B = -CH_2 - C(R^1R^3) - , C = -CH_2 - C(R^1R^4) - , wherein$$

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R¹ is hydrogen or methyl;

R² is -CON(Me)₂, -CO₂CH₂CH₂OH, -OCOCH₃, -OCOCH₂CH₂Ph, -OH or a lactam group;

R³ is -CON(Me)₂, -CO₂CH₂CH₂OH, -OCOCH₃, -OCOCH₂CH₂Ph, -OH or a lactarn group when B is -CH₂-C(R¹R³)- with the proviso that R² and R³ are not the same unless R^2 and R^3 is -OH; and

R⁴ is -R⁵C(O)P(O) R⁶R⁷ or -R⁵P(O)R⁶OC(O)R⁷, wherein R⁵, R⁶ and R⁷ are selected among same or different aryl groups comprising phenyl, methylphenyl, dimethylphenyl, trimethylphenyl, methoxyphenyl, dimethoxyphenyl, trimethoxyphenyl, methylolphenyl, dimethylolphenyl, trimethylolphenyl or styryl radicals.

- 6. Photocrosslinkers according to claim 5, wherein R² and R³ are selected so as to form a water-soluble molecule.
- 7. Photocrosslinkers according to claim 5, wherein said lactam units together with units A or B constitute N-vinylpyrrolidone units.
- 8. Photocrosslinkers according to claim 5, wherein at least one of R² and R³ is hydroxyl.
- 9. Photocrosslinkers according to claim 5, wherein A is N-vinylpyrrolidone, B is vinyl alcohol.
- 10. Photocrosslinkers according to claim 1 or 5 provided with functional groups for crosslinking.
- 11. Photocrosslinkers according to claim 10 provided with functional groups selected among vinylic, acrylic and methacrylic groups.
- 12. Photocrosslinkers according to claim 1 characterized in that units A. B and C are

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siloxane monomer units of a general formula -RaRbSiO-, wherein Ra and Rb in units A and B are selected among lower substituted or unsubstituted alkyl groups, aryl groups and arylalkyl groups.

- 13. Photocrosslinkers according to claim 12, wherein at least on of Ra and Rb is an aryl or arylalkylgroup.
- 14. Photocrosslinkers according to claim 13, wherein at least one of R_a and R_b is substituted with one or more fluorine atoms.
- 15. Photocrosslinkers according to claim 1, wherein units A, B, C are siloxane units comprising substituents in accordance with:

A is $-Si(R^1R^2)$ -O-, B is $-Si(R^1R^3)$ -O- and C is $-Si(R^1R^4)$ -O-, wherein

R¹ is C1 to C6 alkyl; R² is C1 to C6 alkyl or phenyl; R³ is R¹, R² or C1 to C6 fluroalkyl;

R⁴ is -R⁵R⁶C(O)P(O) R⁷R⁸ or -R⁵R⁶P(O)R⁷OC(O)R⁸, wherein R⁵ is a spacing group; R⁶, R⁷ and R⁸ are selected among same or different aryl groups comprising phenyl, methylphenyl, dimethylphenyl, trimethylphenyl, methoxyphenyl, dimethoxyphenyl, trimethoxyphenyl, methylolphenyl, dimethylolphenyl, trimethylolphenyl or styryl radicals.

- 16. Photocrosslinkers according to claim 15, wherein R⁵ is aliphatic spacing group comprising between one and ten atoms.
- 17. Photocrosslinker according to claim 16, wherein said spacing group is (-CH₂)_n, wherein n is between 1 and 10.
- 18. Photocrosslinkers according to claim 15, wherein R1 is methyl; R2 is methyl or phenyl; R³ is R¹. R² or -CH₂CH₂CF₃.

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- 19. Photocrosslinkers according to claim 15 having functional acrylic groups in its terminal ends.
- 20. A method of forming a macromolecular crosslinked network from a composition comprising a photocrosslinker according to any of claims 1 to 19 by irradiating said composition with light exceeding a wavelength of about 305 nm for a time sufficient to form a solid article.
- 21. A method forming a macromolecular crosslinked network from a composition comprising a photocrosslinker according to any of claims 1 to 11 and at least one copolymerizable vinylic, acrylic or methacrylic monomer.
- 22. A method according to claim 20, wherein said composition further comprises a polymer provided with functional vinylic, acrylic or methacrylic groups.
- 23. A method according to claim 22, wherein said polymer has a backbone of ethylene units.
- 24. A method according to claim 22, wherein said polymer is a polysiloxane.
- 25. A method according to any of claims 20 to 24, wherein an ophthalmic lens is produced.
- 26. A method according to claim 25, wherein an intraocular lens is produced in the capsular bag of the eye.
- 27. An ophthalmically acceptable composition comprising photocrosslinkers according to any of claims 1 to 19, having a refractive index greater than about 1.39 and a viscosity such that said composition can be injected through standard cannula having a needle of 15 Gauge, or finer.

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28. The use of photocrosslinkers according to any of claims 1 to 19 in an ophthalmologically acceptable composition for injection into the capsular bag of the eye.

DECLARATION and POWER OF ATTORNEY

U.S. NATIONAL PHASE OF INTERNATIONAL APPLICATION

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name.

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled Macromolecular Compounds, the specification of which was filed as International Application No. PCT/EP00/02539 on March 16, 2000.

ГТ	and was amended under Article 19 on	
LJ		(if applicable)
[]	and was amended under Article 34 on	·
		(if applicable)
[X]	and was assigned U.S. Application Serial No.	09/936,647, and was

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above.

amended on September 14, 2001.

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, §1.56(a).

I hereby claim foreign priority benefits and/or U.S. Provisional application priority benefits under Title 35, United States Code, §119 of any foreign application(s) or U.S. Provisional applications for patent or inventor's certificate listed below and have also identified below any foreign application for patent or inventor's certificate having a filing date before that of the application on which priority is claimed:

			Priority	Claimed
Number	Country	Day/Month/Year Filed	Yes	No
9900935-9	Sweden	March 16, 1999	X	

I hereby claim the benefit under Title 35, United States Code, §120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code, §112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulation, §1.56(a) which

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occurred between the filing date of the prior application and the PCT international filing date of this application:

(Application Serial No.) (Filing Date) (Status) (patented, pending, abandoned)

I hereby appoint Holly D. Kozlowski, Registration No. 30,468; Ronald J. Snyder, Registration No. 31,062; James D. Liles, Registration No. 28,320; Lynda E. Roesch, Registration No. 29,696; Martin J. Miller, Registration No. 35,953; John V. Harmeyer, Registration No. 41,815; Scott N. Barker, Registration No. 42,292; Stephen S. Wentsler, Registration No. 46,403, and Ryan O. White, Registration No. 45,541; Charles H. Brown III, Registration No. 48,866; Jeffrey R. Schaefer, Registration No. 48,514; Todd W. Minor, Registration No. 48,965; John F. Colligan, Registration No. 48,240; Rebecca A. Brown, Registration No. 47,452; and Clayton R. Kuhnell, Registration No. 48,691, my attorneys, c/o Dinsmore & Shohl, 1900 Chemed Center, 255 East Fifth Street, Cincinnati, Ohio 45202 (513) 977-8200, my attorneys, with full power in each of them, of substitution and revocation, to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith.

The undersigned hereby authorizes the above-named U.S. attorneys to accept and follow instructions from **Pharmacia Groningen BV** as to any action to be taken in the Patent and Trademark Office regarding this application without direct communication between the undersigned and the aforenamed U.S. attorneys. In the event of a change in the firm or persons from whom instructions may be taken, the aforenamed U.S. attorneys will be so notified in writing by the undersigned.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

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